ClinVar

NCBI's ClinVar is a freely available submission-driven database for information about genomic variation and its relationship to human health.



ClinVar accepts submissions interpretations of genetic data from:

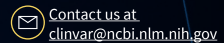
- clinical genetics testing laboratories
- research groups
- expert panels
- and others

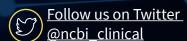


Interpret your data and guide your diagnosis

ncbi.nlm.nih.gov/clinvar

- 1,670+ submitters
- 75+ countries
- 841,000+ variants
- 1,300,000+ submitted records
- ClinVar Search Video





Visit us at ncbi.nlm.nih.gov/ clinvar/ to find out more



ClinVar aggregates





ClinVar helps clinicians

interpret genetic test results

and diagnose disorders to

improve patient outcomes.

What's New

<u>Automated validation</u> in the <u>ClinVar Submission Portal</u> for quick resolution of common errors and faster submission processing time

Tailored <u>notifications</u> for changes in clinical interpretation of variants

Submit to ClinVar

Submit Now

- Setup and register Go to <u>ClinVar Submission Portal</u> to create your myncbi account and register your organization
- **Submit** Use the submission wizard for a single variant submission or excel, TSV/CSV, or XML formats for multiple submissions
- Review and access –. Your data will be available on <u>ClinVar</u> after curatorial review and processing

Dowload data from the ClinVar FTP Site

dbGaP



An NIH-sponsored repository for archiving, curating, and distributing information produced by genome-scale studies investigating the interaction of human genotype and phenotype

Augment your research

View Map

Over

2.6 million

research subjects

Over

1,500 research studies

Over

350,000 variables

Over

100,000

samples of non-genomics omics data

Over

400,000

whole genome and whole exome sequences related to dbGaP studies, available on Amazon Web Services and Google Cloud

dbGaP study submission steps (NIH funded studies)

Registration

- Contact NIH Program Officer or Genomic Program Administrator (GPA)
- Receive invitation
- · Enter study metadata

2 Submission

- Use dbGaP submission guide to upload files
- Work with curators to complete submission
- · Get accession number

3 Release

- · Approve processed data
- Release study

Submit Now

Upcoming

- Public API for study metadata and controlled-access data access using <u>FHIR</u> (Fast Healthcare Interoperability Resources) protocol
- Automated validation in <u>dbGaP Submission Portal</u> for quick feedback and shorter submission processing timeframes

dbGaP study access steps (for Principal Investigators (PIs)

1 Account Setup

- NIH Intramural researchers

 submit permission form to
 establish data request eligibility
 in dbGaP
- Other researchers Get eRA commons user account

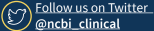
2 Access Application

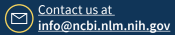
- Complete / revise and submit application to Signing Officer (SO)
- SO certifies application with one or more Data Access Requests (DAR)

3 Approval and Access

- dbGaP Data Access Committee (DAC) reviews and approves application
- dbGaP approved data is provided for download









NIH Genetic Testing Registry (GTR®)



An unbiased, free to participate in, and free to use, international database of clinical and research molecular, cytogenetics and biochemical genetic tests, and supporting information

ABOUT GTR®

- Single gene tests, panels, genomes, and exomes
- 76,000+ tests (incl. 1,600+ tests for somatic targets), 16,000+ conditions, 18,500+ genes, 575+ labs
- 56 BRCA1 single gene tests and 391 multigene panels
- 78 BRCA2 single gene tests and 445 multigene panels

GTR® now includes molecular and serological tests for microbes that affect human health & disease

- 18 COVID-19 tests
- 15 tests for other viruses, parasites, bacteria

GTR® is a central location for laboratories to provide genetic test information and for clinicians and researchers to search and find genetic tests. GTR® increases transparency in the genetic testing landscape.



Submit to GTR®

- Get access and register your lab -Review the **Submission Portal Guide** for details on how to create your account and submit information
- Submit Provide your test data via online forms and/or excel template for bulk submissions
- Review and access GTR processes your tests quickly, assigns it a unique identifier (GTR_ID) and makes it public for users to review

Submit

Starting October 2020, registration of microbe tests opens, so you can find:

- molecular tests to detect microbe nucleic acids
- tests to detect microbe-specific antigens
- · tests to detect antibodies to a microbe
- microbe panels
- · viral load tests to monitor disease progression and guide treatment

LEARN ABOUT GENETIC TESTS AVAILABLE TO YOU

Visit GTR



Purpose and limitations



Clinical utility



Methodology



Clinical and analytical validity



Lab contacts and credentials. including CLIA and state licenses



AMA CPT® and LOINC codes



Evidence of the test's usefulness

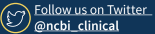


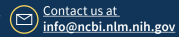
Test ordering information



Test targets (e.g. genes, variants, proteins)

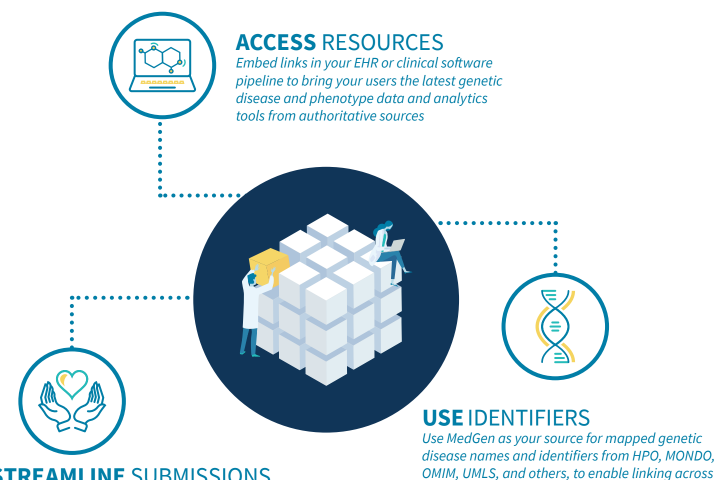












STREAMLINE SUBMISSIONS

MedGen is the phenotype backbone of ClinVar and GTR. Facilitate your organization's submissions by using the disease identifiers in MedGen

RESOURCES

MedGen supports research, diagnosis and treatment of genetic disorders by providing information on:

- Mendelian disorders
- Pharmacogenetic responses
- Complex diseases
- Clinical findings

TOOLS

MedGen's all-in-one platform connects clinicians to leading genetic resources, including:

- PubMed
- GARD
- GeneReviews®

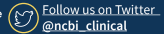
resources

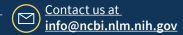
OMIM



Visit MedGen









MedGen, ClinVar, and GTR®

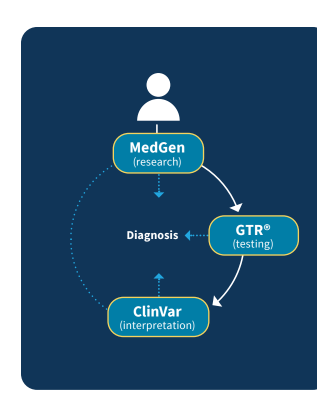


Using NCBI's medical genetics and human variation resources to research, diagnose and treat genetic conditions.

CASE STUDY: SUSPECTED MARFAN SYNDROME

A 9-year-old boy comes into the clinic for a medical release to play soccer. The child presents with myopia, arachnodactyly and has a family history of aortic dissection. These are flags that prompt you to investigate a possible genetic disease.

- Search <u>MedGen</u> to help develop a differentby patient's clinical features in ial diagnosis and learn about a condition, its diagnosis and etiology
- Search the NIH <u>Genetic Testing Registry (GTR®)</u> to find the most appropriate genetic test for your patient for example a panel that includes all the conditions in your differential diagnosis. Learn about the test's validity and utility, and find the ordering information
- Following testing, compare the test results to interpreted variants in <u>ClinVar</u> and determine pathogenicity
- Return to MedGen for practice guidelines from medical and professional societies and the latest research to guide your treatment options. Access consumer resources to help your patient and his family understand his diagnosis, prognosis and available help. If the family is interested, check if there are available studies in ClinicalTrials.gov



ABOUT OUR RESOURCES

ClinVar has more than 1,300,000 submitted records representing more than 841,000 unique variants from 1670+ submitters.

GTR® has 76,000+ tests for 16,000+ conditions and 18,500+ genes, from 575+ labs. It includes clinical and research molecular, cytogenetic and biochemical genetic tests.

MedGen helps research thousands of genetic phenotypes including Mendelian disorders, complex diseases, clinical features and drug responses. It aggregates information from authoritative resources so from one website you can access most available clinical, consumer and molecular resources.

Share Your Data with ClinVar

Share Your Data with GTR



